

1346098

PATENT SPECIFICATION

(11) **1346098**

(21) Application No. 33026/71 (22) Filed 14 July 1971
(31) Convention Application No. 62278 (32) Filed 14 July 1970
(31) Convention Application No. 29012 (32) Filed 1 May 1971
(31) Convention Application No. 33818 (32) Filed 18 May 1971 in
(33) Japan (JA)
(44) Complete Specification published 6 Feb. 1974
(51) International Classification C08F 29/38, 37/00 // D06M 15/36
(52) Index at acceptance



C3P 11C13C 11C14A 11C17 11D2A2A 11K8 2C13B 2C13C
2C14A 2C17 2D1A1 2K8 8C13A 8C13B 8C13C
8C14A 8C17 8C18 8C3 8D2A 8D2B2 8D4 8D8
8K11 8K2 8K8
B2E 179 180 182 186 189 18Y 207 216 21Y 227 228
236 239 23Y 246 24Y 263 275 286 289 28Y 296
299 29Y 30X 319 327 32X 32Y 339 389 410 44Y
469 46Y 473 483 485 486 487 497 498 49Y 505
507 50X 50Y 510 513 51X 532 533 534 535 536
543 544 545 546 547 54Y 555 55Y 560 565
567 568 56Y 575 576 577 583 586 587 588 58Y
C3R 35C11 35C16 35C29 35C8P 35L2A 35L6G
D1P 21Y 235 23Y 242 24Y 260 262 26X 26Y 270 272
274 27Y 361 365 36Y 370 580 587 588 595 59Y
610 612 632 634 70Y 710 71Y 720 722 72Y 73X
73Y 743 744 74X 74Y 760 770 77Y L2 L6

(54) WATER- AND OIL-REPELLENT COMPOSITIONS

(71) We, DAIKIN KOGYO KABUSHIKI KAISHA, a body corporate organised and existing under the Laws of Japan, of Shin-hankyu Bldg., 8 Umeda, Kita-ku, 5 Osaka-shi, Japan, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—
10 This invention relates to an oil- and water-repellent composition, and more particularly to the improvement of fluorochemical compositions for imparting oil and water repellency to fibrous materials.
15 It is well known in the art that the surface of fibrous materials treated with a certain kind of fluorine-containing polymers displays oil and water repellency. Such fluorine-containing polymers having the above property are those having side chains containing a perfluoroalkyl group of 3 to 21 carbon atoms.
20 However, the fibrous materials treated with the fluorine-containing polymers have the undesirable property of accumulating electrostatic charges developed by friction, which renders them susceptible to soiling. To overcome this disadvantage attempts have been made to use antistatic agents in combination with the fluorine-containing polymers, but the water repellency of the fibrous materials treated therewith decreases due to the hydrophilic property of the antistatic agents.
25 An object of this invention is to obviate or mitigate the aforesaid disadvantages.

The inventors have studied the improvement of a composition containing fluorine-containing polymer and antistatic agent by adding numerous compounds of various kinds to the composition. As a result it has been found that when a water-soluble salt such as ammonium chloride is added to the composition fibrous materials treated with the resultant composition display excellent oil- and water-repellent properties and high order of antistatic property. According to this method, however, it is necessary to use a large amount of antistatic agent if a high order of antistatic property is to be obtained, thus impairing the color fastness to rubbing of fibrous materials dyed with a disperse dye and thereby limiting the application of the composition to a narrow range of uses.

According to the present invention there is provided a water- and oil-repellent composition which comprises an aqueous medium having dispersed therein, in a concentration of 0.05 to 10 weight percent, a fluorine-containing polymer having side chains containing a perfluoroalkyl group of 3 to 21 carbon atoms, a water-soluble salt of guanidine in an amount of 0.001 to 10 times the weight of said fluorine-containing polymer and an antistatic agent in an amount of 0.0003 to 20 times the weight of said fluorine-containing polymer.

According to the present invention it is essential to add a water-soluble guanidine salt in combination with an antistatic agent,

35

40

45

50

55

60

65

[Price 25p]

which enables the fibrous materials treated with the present composition to display high order of oil and water repellency and anti-static property free of deterioration in color fastness to rubbing of the fibrous materials dyed with disperse dyes. Such outstanding effect cannot be obtained with water-soluble salts other than the guanidine salts; the use of the other salts lowers color fastness to rubbing of the dyed fibrous materials.

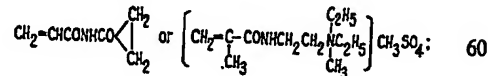
The guanidine salts used in the invention are water-soluble guanidine salts of various acids. The acids include 1) inorganic mono- or poly-basic acids such as hydrochloric acid, nitric acid, carbonic acid, sulfuric acid, phosphoric acid, etc.; 2) organic monobasic acids such as acetic acid, formic acid, gluconic acid, lactic acid, etc. 3) organic polybasic acids such as oxalic acid, malonic acid, succinic acid, malic acid, maleic acid, methylmalonic acid, ethylmalonic acid, tartaric acid, vinaconic acid, glutaric acid, glutaconic acid, saccharic acid, itaconic acid, adipic acid, phthalic acid, dimer of acrylic acid and the like dibasic acids, citric acid, aconitic acid, tricarballylic acid, campholonic acid, trimer of acrylic acid trimer of maleic acid and the like tribasic acids, and polyacrylic acid of a polymerization degree of 4 to 10000, poly-maleic acid of a polymerization degree of 4 to 10000, etc. Of the guanidine salts of these acids the most preferable are guanidine salts of organic monobasic acids and organic poly-basic acids, since such salts ensure more excellent antistatic property than guanidine salts of inorganic acids. The guanidine salts of these acids are easily available and can be prepared in conventional manner, for example by reacting guanidine with an equivalent or excess amount of acid in the presence of a liquid medium such as water or a mixture of water and water-soluble organic solvents. The reaction can be conducted at room temperature or a moderately-elevated temperature of less than 100°C. When polybasic acid is used the reaction product contains minor amounts of byproducts, but there is no need to separate such byproducts because the presence thereof in mixture with the guanidine salts gives no adverse affect on the properties of the present composition.

The antistatic agents used in combination with the above guanidine salts include those which have been used previously as antistatic agents for fabric materials. Examples thereof are as follows:

(1) polymers of acrylamide derivatives

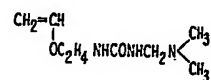
- 95 a) $R^1O(CH_2CH_2O)_nOCCR^2=CH_2$
- b) $CH_2=CR^2COO(CH_2CH_2O)_nOCCR^2=CH_2$
- c) $R^1O[CH(CH_3)CH_2O]_nOCCR^2=CH_2$
- d) $CH_2=CR^2COO[CH(CH_3)CH_2O]_nOCCR^2=CH_2$
- e) $R^1O(CH_2CH_2O)_n[CH(CH_3)CH_2O]_mOCCR^2=CH_2$
- f) $CH_2=CR^2COO(CH_2CH_2O)_n[CH(CH_3)CH_2O]_mOCCR^2=CH_2$

such as a polymer of a compound having the formula of



2) polymers of acrylic ester derivatives such as a polymer of diethylaminoethyl methacrylate quaternized with dimethyl sulfate;

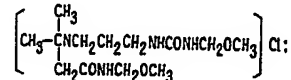
3) polymers of vinyl ethers such as a 65 polymer of a compound having a formula of



4) polymers of vinyl pyridine derivatives such as poly - 2 - vinyl pyridine quaternized with p-toluene sulfonic acid; 70

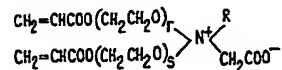
5) polyamine resins such as polyethylene polyamine obtained by reacting a poly-amine with a reaction product of polyethylene glycol and epichlorohydrin;

6) cationic resins such as a polymer of a compound having the formula of



7) amide-based antistatic agents such as 80 N-methoxymethyl polyamide, N,N' - dimethyl - N,N' - bis(hydroxymethyl) adipic amide;

8) amphoteric external permanent antistatic agents such as a compound having the formula of 85

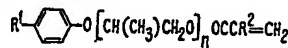
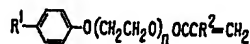


wherein R is alkyl and r and s are each integers from 1 to 100; and

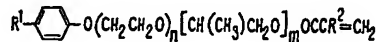
9) polymers containing in the molecule alkylene glycol chains having the formula $-CH_2CH_2-O-$ or $-CH(CH_3)CH_2O-$, 90 such as polymers of the following monomers a) to q) below.

g)

h)



i)

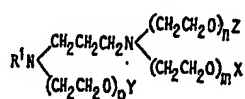
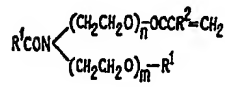


5

j) $\text{R}'\text{COO}(\text{CH}_2\text{CH}_2\text{O})_n\text{OCCR}^2=\text{CH}_2$
 k) $\text{R}'\text{COO}[\text{CH}(\text{CH}_3)\text{CH}_2\text{O}]_n\text{OCCR}^2=\text{CH}_2$
 l) $\text{R}'\text{COO}(\text{CH}_2\text{CH}_2\text{O})_n[\text{CH}(\text{CH}_3)\text{CH}_2\text{O}]_m\text{OCCR}^2=\text{CH}_2$

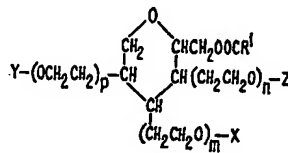
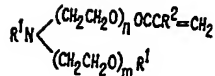
m)

o)



n)

p)



10 wherein R' is hydrogen or an alkyl group of 1 to 20 carbon atoms and R^2 is hydrogen or methyl. At least one of X , Y and Z represents $-\text{OCCR}^2=\text{CH}_2$, $-\text{CH}=\text{CH}_2$ or $-\text{CCl}=\text{CH}_2$ and any remainder are R' or $\text{R}'\text{CO}$, R' and R^2 being as defined above. Independently, n , m and p are a number of from 1 to 100, and q is a number of from 1 to 200.

15 Of these antistatic agents the polymers 9) which contain alkylene glycol chain in the molecule are the most preferable since they ensure the most preferable effects when used in combination with the guanidine salts disclosed before. Such polymers 9) include homopolymers of the monomers a) to q) above, 20 copolymers of these monomers with one another and copolymers of such monomers a) to q) with other copolymerizable monomers containing the copolymerised monomers a) to q) in an amount of at least 70 weight percent. 25 Examples of the copolymerizable monomers are (1) acrylic acid, methacrylic acid or such esters thereof as methyl, ethyl, propyl, butyl, isobutyl, 2-ethylhexyl, hexyl, decyl, lauryl, 30

stearyl, etc.; (2) vinyl esters of aliphatic acids, such as vinyl acetate, vinyl propionate, vinyl caprylate, vinyl laurate, vinyl stearate, etc.; (3) styrene or styrene compounds, such as α -methylstyrene, p -methylstyrene, etc.; (4) halogenated vinyl or vinylidene compounds such as vinyl fluoride, vinyl chloride, vinyl bromide, vinylidene fluoride, vinylidene chloride, etc.; (5) allyl esters of aliphatic acids, such as allyl heptanoate, allyl caproate, allyl caprylate, etc.; (6) vinylalkyl ketones, such as, vinylmethyl ketone, vinylethyl ketone, etc.; (7) acrylamides, such as N -methylol acrylamide, N -methylol methacrylamide, glycidyl acrylate, glycidyl methacrylate, etc.; and (8) diene compounds, such as butadiene, 2,3-dichloro-1,3-butadiene, isoprene, etc. Polymerization reaction for producing these polymers may be carried out by the method conventional to the polymerization for various vinyl monomers. According to one of preferred methods, for instance, the monomer or monomers are polymerized with stirring in organic solvents to produce the desired polymers.

35

40

45

50

55

Fluorine-containing polymers used in the invention are those having side chains containing a perfluoroalkyl group of 3 to 21 carbon atoms, and examples of such polymers 5 are homopolymers or copolymers of the following monomers:

No.	Formula
	R^3
(1)	$R_2SO_2NR^4OOCR^5=CH_2$
(2)	$R_2(CH_2)_nOOCR^5=CH_2$
	R^3
10	(3) $R_2CONR^4OOCR^5=CH_2$ OH
	(4) $R_2CH_2CHCH_2OOCR^5=CH_2$ OOCR ⁶
	(5) $R_2CH_2CHCH_2OOCR^5=CH_2$
	(6) $R_2CH_2COOCH=CH_2$
	(7) $R_2CH=CH(CH_2)_nOOCR^5=CH_2$

15 wherein R_2 is a perfluoroalkyl group of 3 to 21 carbon atoms, R^3 is hydrogen or an alkyl group of 1 to 10 carbon atoms, R^4 is an alkylene group of 1 to 10 carbon atoms, R^5 is hydrogen or methyl, R^6 is an alkyl group of 20 1 to 17 carbon atoms and n is an integer of 1 to 10.

25 These monomers and polymers thereof are known in the art. For example, polymers of the monomer (1) are disclosed in British Patent No. 857,689 and No. 904,262, those of the monomer (2) are in U.S. Patent No. 2,642,416 and No. 3,102,103 and British Patent No. 1,011,612, those of the monomer (3) are in U.S. Patent No. 2,764,603, those of the monomers (4) are in British Patent No. 1,095,900, those of the monomers (5) are in British Patent No. 1,123,379, those of the monomer (6) are in U.S. Patent No. 2,592,609 and those of the monomer (7) are 35 in British Patent No. 1,101,049. Of these polymers homopolymers or copolymers of the monomers (4) and (5) are most desirable in the invention.

40 Copolymers of the above fluorine-containing monomers include those prepared by copolymerizing the above monomers with one another or by copolymerizing the above monomers with copolymerizable monomers of other kinds. Examples of the copolymerizable monomers of other kinds are (1) acrylic acid, methacrylic acid or such esters thereof as methyl, ethyl, propyl, butyl, isobutyl, 2-ethylhexyl, hexyl, decyl, lauryl, stearyl, etc.; (2) vinyl esters of aliphatic acids, such as 45 vinyl acetate, vinyl propionate, vinyl caprylate, vinyl laurate, vinyl stearate, etc.; (3) styrene or styrene compounds, such as α -methylstyrene, p-methylstyrene, etc.; (4) halo-

50 genated vinyl or vinylidene compounds such as vinyl fluoride, vinyl chloride, vinyl bromide, vinylidene fluoride, vinylidene chloride, etc.; (5) allyl esters of aliphatic acids, such as allyl heptanoate, allyl caproate, allyl caprylate, etc.; (6) vinylalkyl ketones, such as, vinylmethyl ketone, vinylethyl ketone, etc.; (7) acrylamides, such as N-methylol acrylamide, methylol methacrylamide, glycidyl acrylate, glycidyl methacrylate, etc.; and (8) diene compounds, such as butadiene, 2,3-dichloro-1,3-butadiene, isoprene, etc. Of these comonomers acrylic acid, methacrylic acid and esters thereof are most desirable. The copolymers used in the invention contain the copolymerised fluorine-containing monomers in an amount at least 25 weight percent, preferably more than 50 weight percent.

55 Polymerization of the fluorine-containing monomers may be carried out by the methods known to the art, for example, by bulk polymerization, emulsion polymerization and suspension polymerization, using heat, free-radical catalysts, ultraviolet light or ionizing radiations. Of these methods, the most desirable is emulsion polymerization, since stable aqueous polymer emulsion can be obtained by such a method. The emulsion polymerization may be carried out, for example, by polymerizing fluorine-containing monomers in an aqueous medium containing dispersant and free radical initiator. As the dispersant various cationic or anionic surfactants may be used. Examples thereof are octadecyltrimethyl ammonium acetate, dodecyltrimethyl ammonium bromide, tetradodecyltrimethyl ammonium chloride, (dodecylmethyl-benzyl)-trimethyl ammonium chloride, (dodecyl-benzyl)-trimethyl ammonium bromide, pentadecafluorooctyl ammonium chloride and like cationic surfactants, and sodium alkyl ($C_{12}-C_{18}$) benzene sulfonate, sodium alkyl naphthalene sulfonate, sodium oleate, sodium alkyl ($C_{16}-C_{18}$) sulphonate, ammonium perfluoroalkanionate and like anionic surfactants.

60 The aqueous oil- and water-repellent composition of the invention may be prepared easily by dissolving water-soluble guanidine salts in combination with antistatic agent in aqueous polymer emulsion containing fluorine-containing polymers obtained by emulsion polymerization of the fluorine-containing monomers aforementioned or obtained by dispersing fluorine-containing polymers in water. The composition contains the fluorine-containing polymer in an amount of 0.05 to 10 weight percent, preferably 0.3 to 5 weight percent, based on the total weight of the composition. The water-soluble guanidine salt is contained in the composition in an amount of 0.001 to 10 times, preferably 0.5 to 5 times, the weight of the fluorine-containing polymer. Smaller amounts of the salts result in no or 65 poor improvement of water repellency, while larger amounts adversely affect the stability.

of the resultant composition without any better effects. 30

5 The antistatic agents are added to the composition in an amount of 0.0003 to 20 times, preferably 0.5 to 10 times, the weight of the fluorine-containing polymer. 35

10 The present water- and oil-repellent composition preferably has a total solids concentration of not higher than 50 weight percent. 40

15 Fibrous materials which can be treated with the present composition are, for example, yarns, textiles, knitted fabrics, felts and non-woven fabrics made of various fibers, leather 45 and paper. Such various fibers include cellulose, silk, wool and like natural fibers; cellulose acetate, cellulose propionate and like artificial fibers, and polyamide, polyester, polyacrylonitrile, polyvinylalcohol and like synthetic fibers. 50

20 The fibrous materials are wetted with the present composition by coating, dipping, spraying, padding and like conventional methods, so as to attach the fluorine-containing polymer to the fibrous materials in an amount of 0.01 to 5 weight percent in terms of dry weight, based on the weight of the fabric. The wetted fibrous materials are then dried, and cured as required. After curing the 55

25

(CF₃)₂CF(CF₂)₆CH₂—CH(OH)CH₂OOCCH=CH₂

30 N-methylolacrylamide and 2-ethylhexyl methacrylate. 60

35 In a 300-ml 4-necked flask equipped with a reflux condenser, thermometer, dropping funnel and agitator were placed 36 grams of

40 (CH₃)₂CF(CF₂)₆CH₂CH(OH)CH₂OOCCH=CH₂, 65

45 0.34 gram of N-methylolacrylamide, 31.5 grams of 2-ethylhexyl methacrylate and 45 grams of deoxygenated water, 7 grams of acetone and 6.4 grams of a 62:38 weight ratio mixture of dimethyloctadecyl amine and glacial acetic acid. The resultant mixture was 70 thoroughly stirred in nitrogen stream and gradually heated to 40—55°C. To the mixture was added dropwise 0.06 gram of azobisisobutyroamidine hydrochloric acid salt dissolved in 5 grams of deoxygenated water. 75 After the addition the reaction system was heated with stirring at 58—62°C for 3 hours whereby stable polymer emulsion, milky white in colour having a polymer concentration of 50% was obtained. 80

(b) Preparation of water- and oil-repellent composition and treatment of fibrous materials. 85

100 Parts of the resultant polymer emulsion was mixed with aqueous solution of 20 parts of "TAKENON-AS 100" (trade mark, antistatic agent containing a polymer of polyethylene glycol methacrylate, Shin-Nakamura

90 Kagaku Kogyo Kabushiki Kaisha, Japan) dissolved in 2000 parts of water. To the resultant mixture was added with stirring 10 parts of guanidine hydrochloride dissolved in 2000 parts of water and the mixture was diluted with 10,000 parts of water to produce water- and oil-repellent composition, milky white in color. 95

95 15 parts of "Amundsen" fabric (a crepe weave fabric) of polyester was dipped in the resultant composition for 3 minutes and squeezed to 80 percent impregnation, based on the weight of the fabric treated. Then the fabric was dried at 80°C for 20 minutes and heat-treated at 150°C for 3 minutes. 100

100 The fabric thus treated showed no change in color and softness and displayed the following properties. 105

105 Water repellency: 100
Oil repellency: 7
Electrostatic voltage and half-value period: 7V, 1sec. (measured by using Nylon taffeta fabric rubbed at 600 r.p.m. under 110

a load of 500 g at 20°C at 65% RH for 3 minutes, the measurement being made in the same conditions as above herein below)

5 No change was observed in these properties after washing or dry-cleaning the fabric 3 times.

A crêpe weave fabric, known as "Amundsen fabric", of polyester dyed with "Sumikaron Blue R" (trade mark, disperse dye of Sumitomo Chemical Co. Ltd., Japan) was treated with the composition obtained above in the same manner as above, whereby fabric having the following properties was obtained. 10

	Fabric obtained	Contrast (non-treated dyed fabric)
Water repellency:	100	0
Oil repellency:	7	0
Electrostatic voltage and half-value period:	9V, 1 sec.	460 V, 6 sec.
Color fastness:	<div style="display: flex; align-items: center; justify-content: space-between;"> <div style="flex: 1; text-align: center;"> 4—5 degrees in dry state </div> <div style="flex: 1; text-align: center;"> 4—5 degrees in dry state </div> </div>	<div style="display: flex; align-items: center; justify-content: space-between;"> <div style="flex: 1; text-align: center;"> 4—5 degrees in wet state </div> <div style="flex: 1; text-align: center;"> 4—5 degrees in wet state </div> </div>

20 For comparison the same dyed fabric was treated in the same manner as above using the same composition except that no guanidine hydrochloride was added to the composition, whereby the fabric having the following properties was obtained.

Water repellency: 80+

Oil repellency: 7

Electrostatic voltage and half-value period:

25 250 V, 4 sec.

Colorfastness to rubbing:

4—5 degrees
in dry state

4—5 degrees
in wet state

Color fastness to rubbing: 4—5 degrees in dry state, 4—5 degrees in wet state

30 Further, for comparison, the same dyed fabric was treated in the same manner as above using the same composition except that no antistatic agent was added to the composition, whereby the fabric having the following properties was obtained.

Water repellency: 100

Oil repellency: 7

Electrostatic voltage and half-value period: 1200 V, 11 sec.

Colorfastness to rubbing:

4—5 degrees
in dry state

4—5 degrees
in wet state

Comparative Example 1.

Water- and oil-repellent composition was prepared in the same manner as in Example 1 except that ammonium chloride was used in the amount shown below in the place of the guanidine hydrochloride and the antistatic agent, "TAKENON-AS 100" (trade mark, the same as in Example 1), was used in the amount shown below. 40

The same dyed fabric of polyester as in Example 1 was treated with the above composition in the same manner as in Example 1, whereby fabric having the following properties was obtained. 45 50

Amount of antistatic agent used (parts)	20	40	40								
Amount of NH_4Cl used (parts)	10	10	20								
Water repellency:	100	100	100								
Oil repellency:	7	7	7								
Electrostatic voltage and half-value period:	330 V 8 sec.	110 V 5 sec.	15 V 2 sec.								
Color fastness to rubbing:	<table border="0"> <tr> <td>In dry state</td> <td>4-5</td> <td>3</td> <td>3</td> </tr> <tr> <td>In wet state</td> <td>4-5</td> <td>2-3</td> <td>2-3</td> </tr> </table>	In dry state	4-5	3	3	In wet state	4-5	2-3	2-3		
In dry state	4-5	3	3								
In wet state	4-5	2-3	2-3								

Example 2.

(a) Preparation of copolymer of $(\text{CF}_3)_2\text{CF}(\text{CF}_2)_6\text{CH}_2\text{CHCH}_2\text{OOCCH}=\text{CH}_2$, N-methylolacrylamide and 2-ethylhexyl methacrylate.

Aqueous emulsion having 50% concentration of copolymer of



10 N-methylolacrylamide and 2-ethylhexyl methacrylate was prepared in the same manner as in Example 1—(a) except that



was used in the place of



15 (b) Preparation of water- and oil-repellent composition and treatment of fibrous material.

0.5 part of the resultant aqueous polymer emulsion obtained as above (a) was mixed with stirring with 0.5 part of "TAKENON-AS 100" (trade mark, the same as in Example 1) dissolved in 20 parts of water and 0.5 part of guanidine hydrochloride dissolved in 50 parts of water. The resultant mixture was diluted with water to produce 100 parts of aqueous water- and oil-repellent composition, milky white in color.

The same dyed fabric as Example 1 was treated with the above composition in the same manner as in Example 1, whereby fabric having the following properties was obtained.

Water repellency: 100

Oil repellency: 7

Electrostatic voltage: 10 V

Color fastness to rubbing:

4-5 degrees in dry state
4-5 degrees in wet state

35

Comparative Example 2.

Water- and oil-repellent composition was prepared in the same manner as in Example 2 except that ammonium chloride was used in the amount shown below in the place of guanidine hydrochloride and the antistatic agent was used in the amount shown below.

The same dyed fabric of polyester as Example 1 was treated with the above composition in the same manner as in Example 1, whereby fabric having the following properties was obtained.

40

45

Amount of antistatic agent used (parts)	0.5	1	1
Amount of NH_4Cl used (parts)	0.5	0.5	1
Water repellency:	100	100	100
Oil repellency:	7	7	7
Electrostatic voltage:	320 V	12 V	95 V
Color fastness to rubbing:	<div style="display: flex; align-items: center;"> In dry state 4-5 3 3 </div>		
	<div style="display: flex; align-items: center;"> in wet state 4-5 2 2 </div>		

Example 3.

(a) Preparation of the homopolymer of



5 In the same flask as in Example 1 were placed 100 grams of



10 5 grams of trimethyloctadecyl ammonium bromide, 1 gram of azobisisobutyroamidine hydrochloride, 15 grams of acetone and 80 grams of deoxygenated water, and heated at 58-62°C in nitrogen stream for 6 hours.

The polymer emulsion thus obtained had a polymer concentration of 50%.

15 (b) Preparation of antistatic polymer.

In a 300-ml four-necked flask equipped with a reflux condenser, thermometer, dropping funnel and stirrer were placed 37 g of



20 7 g of glycidyl methacrylate, 200 g of dimethylformamide and 0.3 g of azobisisobutyronitrile. The mixture was stirred homogeneously at room temperature and the temperature of the system was gradually raised to 70°C.

25 The system was stirred at that temperature for 6 hours, whereby slightly viscous polymer solution, light yellow in color, having a concentration of 17% was obtained. Removal of dimethylformamide by distillation gave a viscous polymer. The polymer was yellow in colour and had water-solubility.

30 (c) Preparation of water- and oil-repellent composition and treatment of fibrous material.

1.5 parts of the polymer emulsion obtained as above (a) was mixed with stirring with 0.2 part of antistatic polymer of above (b) dissolved in 20 parts of water and 0.5 part of guanidine hydrochloride dissolved in 20 parts of water. The resultant mixture was diluted with water to produce 100 parts of water- and oil-repellent composition, milky white in color.

35 Taffeta fabric of polyester was treated with the composition obtained as above in the same manner as in Example 1, whereby fabric having the following properties was obtained.

Water repellency:	100	50
Oil repellency:	7	50
Electrostatic voltage:	25 V	

40 For comparison the same fabric as above was treated in the same manner as above except that no antistatic polymer was added to the composition. The fabric thus obtained displayed the following properties.

Water repellency:	100	50
Oil repellency:	7	50
Electrostatic voltage:	2280 V	

45 Further, for comparison, the same fabric as above was treated in the same manner as above except that no guanidine hydrochloride was added to the composition, whereby fabric having the following properties was obtained.

Water repellency:	90	60
Oil repellency:	7	60
Electrostatic voltage:	2350 V	

55 (a) Preparation of the copolymer of



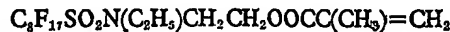
and butadiene.

In a 100-ml pressure ampoule were placed 31.5 grams of deoxygenated water, 4.7 grams of acetone, 1.25 grams of

0.05 gram of potassium persulfate, 0.125 gram of dodecylnercaptan, 0.125 gram of n-octylnercaptan and 12.5 grams of butadiene, and the ampoule was sealed. The ampoule was heated with shaking at 50°C for 26 hours to polymerize the monomers. After this reaction, 12.5 grams of

10

5 $C_8F_{17}SO_2NH(CH_2)_5N(CH_3)_2HCl$,



15 and 8.8 grams of acetone were further added to the reaction mixture and heated at 50°C for 16 hours for further polymerization. Thus, polymer emulsion milky white in color having a polymer concentration of 29 percent was obtained.

20 (b) Preparation of antistatic polymer.

In a 300-ml four-necked flask equipped with a reflux condenser, thermometer, dropping funnel and stirrer were placed 40 g of

$C_{16}H_{35}O(CH_2CH_2O)_{20}OCCH=CH_2$,

25 2 g of N-methylol acrylamide 200 g of dimethylformamide and 0.5 g of azobisisobutyronitrile. The mixture was stirred homogeneously at room temperature and the temperature of the system was gradually raised to 60°C. The system was stirred at that temperature for 10 hours, whereby transparent viscous polymer solution was obtained. The polymer was soluble in acetone and dimethyl formamide but insoluble in water. One part of the resultant polymer solution was mixed with stirring with 99 parts of water to produce 1% aqueous dispersion of the polymer.

30 (c) Preparation of water- and oil-repellent composition and treatment of fibrous material.

40 1.5 parts of the polymer emulsion prepared as above (a) was mixed with stirring with 2.5 parts of 1% aqueous dispersion of antistatic polymer prepared in the above manner (b) and 0.5 part of guanidine hydrochloride dissolved in 20 parts of water. The mixture was diluted with water to produce 100 parts of water- and oil-repellent composition, milky white in color.

45 Tropical fabric of polyester was treated with the above water- and oil-repellent composition in the same manner as in Example 1, whereby fabric having the following properties was obtained.

50 Water repellency: 100
Oil repellency: 7
Electrostatic voltage: 25 V

Example 5.
"Amundsen fabric" dyed with the disperse dyes shown in Table 1 below was treated in the same manner as in Example 1 with the water- and oil-repellent composition obtained in Example 1. The color fastness to rubbing of the resultant fabric in wet state was as follows:

TABLE 1

65

Dyes used	Color fastness to rubbing in wet state (degrees)	
Foron-Blue-E-BL* ¹	4—5	
Foron-Black-E-PW* ²	4	70
Foron-Brown-S-2BL* ³	4	
Foron-Red-E-2GL* ⁴	4	

Note: *1) to *4) are trade marks for disperse dyes of SANDOZ Kabushiki Kaisha, Japan.

75

Example 6.
One part of the polymer emulsion obtained in the same manner as in Example 2—a was mixed with stirring with 0.1 part of "PER-MAX AW-2" (trade mark, antistatic agent containing polymer of quaternary ammonium salt of diethyl aminoethyl methacrylate, Yoshimura Yukagaku Kabushiki Kaisha, Japan) dissolved in 20 parts of water and 0.1 part of guanidine hydrochloride dissolved in 20 parts of water. The resultant mixture was diluted with water to produce 100 parts of water- and oil-repellent composition, milky white in color.

80 2 parts of "Amundsen fabric" of polyester was dipped in the composition thus obtained for one minute and squeezed to 80% impregnation based on the weight of the fabric treated. Then the fabric was dried at 80°C for 10 minutes and heat-treated at 150°C for 3 minutes, whereby fabric having the following properties was obtained.

80

85

90

95

	Fabric treated	Contrast (Non-treated fabric)
Water repellency:	100	0
Oil repellency:	7	0
Electrostatic voltage:	28 V	1900 V

For comparison the same fabric was treated in the same manner as above using the same composition except that guanidine hydrochloride was not added to the composition, which gave fabric having water repellency of 80, oil repellency of 7 and antistatic voltage of 485 V. Further, for comparison, the same fabric was treated in the same manner as above using the same composition except that ammonium chloride was used in the place of guanidine hydrochloride. The fabric thus obtained displayed water repellency of 100 and oil repellency of 7 with high antistatic voltage of 488 V.

Example 7.
One part of the polymer emulsion obtained

in the same manner as in Example 3-a) was mixed with the predetermined amount of "PERMAX AW-2" (trade mark, the same as in Example 6) and guanidine phosphate in the same manner as in Example 6 to produce water- and oil-repellent composition.

"Amundsen fabric" of polyester was treated with the above composition in the same manner as in Example 6. Thus fabric having the properties shown in Table 2 below was obtained. The table also shows the properties of the fabric treated in the same manner as above using the same composition except that ammonium monohydrogen phosphate was used in the place of guanidine phosphate.

20

25

30

TABLE 2

Amount of polymer emulsion (part)	Amount of PERMAX	Amount of guanidine phosphate (part)	Amount of ammonium monohydrogen phosphate (part)	Water repellency	Oil repellency	Electrostatic voltage (V)
1	0.1	0.2	—	100	7	6
1	0.1	—	0.2	100	7	85
1	1.0	0.2	—	100	7	3
1	1.0	—	0.2	100	7	16
1	0.05	0.2	—	100	7	9
1	0.05	—	0.2	100	7	150
1	0.01	0.05	—	100	7	18
1	1.00	1.00	—	100	7	3

Example 8.

(a) Preparation of antistatic polymer.

In a 300-ml four necked flask were placed 5 60 g of dimethyl sulfate salt of 2-dimethylaminoethyl methacrylate and 100 g of methyl Cellosolve. ("CELLIOSOLVE" is a Trade 10 Mark). The mixture was heated at 70°C in nitrogen gas stream and to the mixture was added dropwise 0.05 g of benzoyl peroxide 15 dissolved in 5 g of methyl Cellosolve. After the addition the system was heated with stirring at 70—75°C for 2 hours. To the resultant reaction mixture 35 g of water was added after being cooled, whereby 30% aqueous solution of water-soluble polymer dissolved in a mixture of water and methyl Cellosolve was obtained.

(b) Preparation of water- and oil-repellent composition and treatment of fibrous material.

One part of the polymer dispersion obtained in the same manner as in Example 4—a) was slowly stirred with 0.2 part of 30% aqueous solution of the antistatic agent obtained as above (a) and 0.2 part of guanidine hydrochloride dissolved in 20 parts of water. The resultant mixture was diluted with water to produce 100 parts of water- and oil-repellent composition, milky white in color. 20

Fabric of polyamide was dipped in the

composition thus obtained and squeezed to 25

50% impregnation based on the weight of the fabric treated. Then the fabric was dried at 30

80°C for 10 minutes and heat-treated at 35

140°C for 5 minutes, whereby fabric of the following properties was obtained.

5 Water repellency: 100
Oil repellency: 7
Electrostatic voltage: 20 V

In the same flask as in Example 1-(a) were placed 75 parts of

$C_7F_{15}CH_2OOCCH=CH_2$, 30

Example 9.

10 2 parts of the polymer emulsion obtained in the same manner as in Example 2-a) was mixed with stirring with 1 part of "Aston 900 EP" (trade mark, polyethylene polyamine antistatic agent, Millmaster Onyx Corp., U.S.A.) dissolved in 20 parts of water and 0.2 part of guanidine hydrochloride dissolved in 30 parts of water. The resultant mixture 15 was diluted with water to produce 100 parts of water- and oil-repellent composition, milky white in color.

20 "Amundsen fabric" of polyester, taffeta fabric of polyamide and taffeta fabric of polypropylene were respectively treated with the above composition in the same manner as in Example 1. Each fabric thus treated showed water repellency of 100, oil repellency of 7 and excellent antistatic property.

25 Example 10.

a) Preparation of homopolymer of

$C_7F_{15}CH_2OOCCH=CH_2$.

2.8 parts of sodium lauryl sulfate, 0.4 part of potassium persulfate and 135 parts of de-oxidized water. The mixture was then heated in nitrogen gas stream at 50°C for 3 minutes, whereby polymer emulsion having a concentration of 33% was obtained.

b) Preparation of water- and oil-repellent composition and treatment of fibrous materials.

40 One part of the polymer emulsion obtained above a) was mixed with 0.3 part of "TH-44" (trade mark, aqueous dispersion of paraffin wax and polymer of quaternary ammonium salt of dimethylaminoethyl methacrylate, Nikka Chemical Ind. Co., Ltd., Japan) dissolved in 20 parts of water and 0.2 part of a guanidine salt dissolved in 20 parts of water, the guanidine salt used being specified in Table 3 below. The mixture was diluted with water to produce 100 parts of water- and oil-repellent composition.

45 "Amundsen fabric" of polyester dyed with "Foron Brown S-2BL" (trade mark, the same as in Example 5) was treated with each composition in the same manner as in Example 6, whereby fabric having the properties shown in Table 3 below was obtained.

50

55

55

TABLE 3

Guanidine salt	Water repellency	Oil repellency	Antistatic voltage (V)	Color fastness to rubbing	
				Dry	Wet
Acetate	100	7	18	4 — 5	4 — 5
Sulfate	100	7	22	4 — 5	4 — 5
Nitrate	100	7	30	4 — 5	4 — 5
Phosphate	100	7	18	4 — 5	4 — 5
Non-treated fabric	0	0	580	4 — 5	4 — 5

Example 11.

60 Water- and oil-repellent composition was prepared in the same manner as in Example 8 except that 0.6 part of the polymer emulsion obtained as in Example 2-(a) was used in the place of the polymer emulsion obtained as in Example 4-(a).

65 Taffeta fabric of polyester was dipped in the composition thus obtained for 2 minutes and squeezed at 40% impregnation. Then the fabric was dried at 80°C for 10 minutes and

heat-treated at 150°C for 3 minutes to obtain fabric having water repellency of 100 and oil repellency of 7.

70 The fabric thus treated was forcibly rubbed against taffeta fabric of polyamide sixty times for 30 seconds and brought to a position one cm away from cigarette ashes, but 75 none of the ashes were attracted to the fabric, this showing excellent antistatic property. For comparison non-treated fabric of polyester was rubbed against the taffeta fabric 80

of polyamide as above and brought close to cigarette ashes. Even when the fabric was about 10 cm away from the ashes the fabric attracted a great amount of ashes.

5 Example 12.

(a) Preparation of guanidine malonate.

Aqueous solution of 191 parts of guanidine hydrochloride dissolved in 1500 parts of water was passed through a column filled with 10 1500 parts ion exchange resin (Amberlite IRA-400 [OH] type) ("Amberlite" is a Trade Mark). Then resultant ion exchanged liquid was concentrated to produce 800 parts of guanidine solution. After a part of the 15 guanidine solution was titrated with 0.1 N-hydrochloric acid to determine concentration thereof, the solution was mixed with equivalent amount thereto of 10% aqueous malonic acid solution and the mixture was 20 stirred at room temperature. The resultant mixture was diluted with water to obtain colorless transparent aqueous solution containing 10% reaction product of guanidine and malonic acid.

(b) Preparation of water- and oil-repellent composition and treatment of fibrous material.

0.5 part of polymer emulsion obtained in the same manner as in Example 2-(a) was slowly stirred with 0.5 part of "TAKENON AS-100" (trade mark, the same as in Example 1) dissolved in 20 parts of water and 2 parts of 10% aqueous solution of the reaction product of guanidine and malonic acid obtained as in (a) above. The resultant mixture was diluted with water to produce 100 parts of water- and oil-repellent composition, milky white in color.

30 Tropical fabric of polyester was dipped in the above composition for 3 minutes and squeezed to 70% impregnation based on the weight of the fabric treated. Then the fabric was dried at 80°C for 10 minutes and heat-treated at 150°C for 3 minutes.

35 No change was observed in color and softness of the resultant fabric. The fabric displayed the following properties:

	Fabric obtained	Contrast (Non-treated fabric)
Water repellency:	100	0
Oil repellency:	7	0
Electrostatic voltage:	2 V	500 V

50 For comparison the same fabric was treated in the same manner as above using the same composition except that the reaction product of guanidine and malonic acid was not added to the composition, whereby fabric having the following properties was obtained.

55 Water repellency: 80
Oil repellency: 7
Electrostatic voltage: 350 V

60 Further, for comparison, the same fabric was treated in the same manner as above using the same composition except that "TAKENON AS-100" (trade mark, the same as in Example 1) was not added to the composition. The fabric obtained showed the following properties:

65 Water repellency: 100
Oil repellency: 7
Electrostatic voltage: 410 V

70 Example 13.
Guanidine solution was obtained in the same manner as in Example 12 except that

the ion exchange step was conducted at 50°C. The guanidine solution was mixed with equivalent amount of 10% aqueous maleic acid solution and the mixture was stirred at room temperature. The resulting mixture was diluted with water to produce 10% aqueous solution of reaction product of guanidine and maleic acid.

75 2 parts of the resultant solution was mixed with 1.5 parts of the polymer emulsion obtained in the same manner as in Example 4-(a) and 0.2 part of antistatic polymer dissolved in 20 parts of water, the polymer being obtained in the same manner as in Example 3-(b). The resultant mixture was diluted with water to produce 100 parts of water- and oil-repellent composition, milky white in color.

80 Taffeta fabric of polyester was dipped in the composition and squeezed to 40% impregnation based on the weight of the fabric. Then the fabric was dried at 80°C for 20 minutes and heat-treated at 150°C for 3 minutes, whereby fabric having the following properties was obtained.

95

	Fabric obtained	Contrast (Non-treated fabric)
Water repellency:	100	0
Oil repellency:	7	0
Electrostatic voltage:	8 V	320 V

Example 14.

Twelve kinds of water- and oil-repellent compositions were prepared in the same manner as in Example 12 by using varied amounts, shown in Table 4 below, of 10% aqueous solution of guanidine-malonic acid reaction product or of ammonium chloride, fluorine-containing polymer emulsion (as in Example 12) and "TH-44" (trade mark, the same as in Example 10).

5 "Amundsen fabric" of polyester dyed with "Foron-Blue-E-BL" (trade mark, the same as in Example 5) was treated with the twelve respective compositions in the same manner as in Example 1. The properties of the fabric thus obtained are shown in Table 4 below.

10 15

TABLE 4

Amount of fluorine-containing polymer emulsion (%)	Amount of "TH-44"	Amount of guanidine-malic acid reaction product (%)	Amount of NH ₄ Cl	Water repellency	oil repellency	Electrostatic voltage (V)	Color fastness to rubbing in wet state
0.5	1.0	0.2	—	100	7	6	3 — 4
0.5	1.0	0.05	—	100	7	10	3 — 4
0.5	0.5	0.1	—	100	7	11	4
0.25	0.25	0.05	—	100	7	14	4 — 5
0.15	0.1	0.01	—	100	7	18	4 — 5
0.5	1.0	—	0.2	100	7	88	3
0.5	1.0	—	0.05	100	7	200	3 — 4
0.5	0.5	—	0.1	100	7	115	4
0.25	0.25	—	0.05	100	7	280	4 — 5
0.15	0.1	—	0.01	100	7	330	4 — 5
0.5	1.0	—	—	80	7	380	
Non-treated fabric	—	—	—	0	0	1300	4 — 5

Example 15.

Three kinds of water- and oil-repellent composition were prepared in the same manner as in Example 12 in which the following antistatic agents were respectively used in the place of "TAKENON AS-100" (trade mark, the same as in Example 1).

Antistatic agents used:

"TH-44" (trade mark, the same as in Example 10)

"PERMAX AW-2" (trade mark, the same as in Example 6)

"Aston 900 EP" (trade mark, the same as in Example 9)

15 Tropical fabrics of polyester were treated with respective compositions in the same manner as in Example 12. The fabrics obtained were excellent in properties as in Example 12.

Example 16.

0.5 part of 20% solution of diethylaminoethyl methacrylate - 2 - ethylhexyl methacrylate copolymer dissolved in water-butyl Cellosolve mixture was diluted with 20 parts of water and the solution was mixed with 1.5 parts of 30% polymer emulsion prepared in the similar manner as in Example 1-(a) and 5 parts of 10% solution of guanidine-malonic acid reaction product obtained in the same manner as in Example 12. The resultant mixture was diluted with water to produce 100 parts of water and oil-repellent composition, milky in white in color.

20 "Amundsen fabric" of polyester was treated with the above composition in the same manner as in Example 12, whereby fabric having the following properties was obtained.

20

25

30

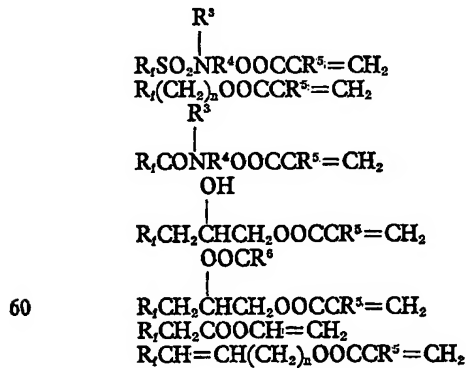
35

	Fabric obtained	Contrast (Non-treated fabric)
Water repellency:	100	0
Oil repellency:	7	0
Electrostatic voltage:	3 V	1380 V

WHAT WE CLAIM IS:—

40 1. A water- and oil-repellent composition which comprises an aqueous medium having dispersed therein, in a concentration of 0.05 to 10 weight percent, a fluorine-containing polymer having side chains containing a perfluoroalkyl group of 3 to 21 carbon atoms, a water-soluble salt of guanidine in an amount of 0.001 to 10 times the weight of said fluorine-containing polymer and an antistatic agent in an amount of 0.0003 to 20 times the weight of said fluorine-containing polymer.

45 2. A composition according to Claim 1, in which said fluorine-containing polymer is a polymer containing at least 25 weight % of units of a monomer selected from monomers having the formulae



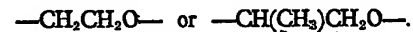
55 wherein R_f is a perfluoroalkyl group of 3 to 21 carbon atoms, R^3 is hydrogen or an alkyl group of 1 to 10 carbon atoms, R^4 is an alkylene group of 1 to 10 carbon atoms, R^5 is hydrogen or methyl, R^6 is an alkyl group of 1 to 17 carbon atoms and n is an integer of 1 to 10.

60 3. A composition according to claim 1 or 2, in which said water-soluble salt of guanidine is a guanidine salt of an inorganic mono- or poly-basic acid.

65 4. A composition according to claim 1 or 2, in which said water-soluble salt of guanidine is a guanidine salt of an organic mono- or poly-basic acid.

70 5. A composition according to claim 1 or 2, in which said water-soluble salt of guanidine is a guanidine salt of an organic poly- basic acid.

75 6. A composition according to any preceding claim, in which said antistatic agent is a polymer containing in the molecule alkylene glycol chains having the formula



80 7. A composition according to claim 6, in which said antistatic agent is a polymer containing at least 70 weight percent of units of a monomer selected from monomers having the formulae

65

70

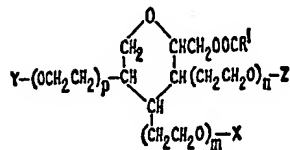
75

80

85

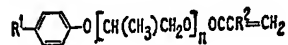
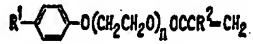
90

5 $R^1O(CH_2CH_2O)_n[CH(CH_3)CH_2O]_mOCCR^2 = CH_2$,
 $CH_2 = CR^2COO(CH_2CH_2O)_nOCCR^2 = CH_2$,
 $R^1O[CH(CH_3)CH_2O]_nOCCR^2 = CH_2$,
 $CH_2 = CR^2COO[CH(CH_3)CH_2O]_nOCCR^2 = CH_2$,

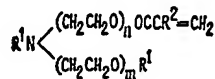
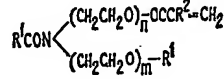


and

$$\text{HS}[\text{CH}_2\text{CHR}^2\text{COO}(\text{CH}_2\text{CH}_2\text{O})_n\text{COCHR}^2\text{CH}_2\text{S}]_q\text{H}$$



10 $R^1COO(CH_2CH_2O)_nOCCR^2=CH_2$,
 $R^1COO[CH(CH_3)CH_2O]_nOCCR^2=CH_2$,
 $R^1COO(CH_2CH_2O)_n[CH(CH_3)CH_2O]_m$
 $OCCR^2=CH_2$



15 

FITZPATRICKS,
Chartered Patent Agents,
14/18 Cadogan Street,
Glasgow, C.2.

and
Warwick House,
Warwick Court,
London, WC1R 5D

has been directed to

2. Subsection (i)

Warwick Square
London, WC1R 5DJ.

he has been directed in

9. Subsection (i) of

to patent No. 1,266,19

Reference has been directed in pursuance of Section 9, Subsection (i) of the Patents Act 1949, to patent No. 1,266,196.

Printed for Her Majesty's Stationery Office by the Courier Press, Leamington Spa, 1974.
Published by the Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from
which copies may be obtained.

THIS PAGE BLANK (USPTO)